PATENT COOPERATION TREATY REC'D 16 FEB 2005

PCT

1	 -000
WIPO	
VVIPO	 PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference 1171WOORD01	FOR FURTHER ACTION	See Form PCT/IPEA/416						
International application No. PCT/EP2004/050376	International filing date (day/more 26.03.2004	nth/year) Priority date (day/month/year) 28.03.2003						
International Patent Classification (IPC) or na	tional classification and IPC							
A61K31/46, A61P11/00, A61K31/44	normal oldoomodilon and it	·						
AURORA, AURORA								
Applicant		•						
ALTANA PHARMA AG et al.								
This report is the international prel Authority under Article 35 and tran	iminary examination report, es smitted to the applicant accor	stablished by this International Preliminary Examining ding to Article 36.						
2. This REPORT consists of a total o	f 6 sheets, including this cove	er sheet.						
3. This report is also accompanied by								
a. \square sent to the applicant and to	the International Bureau) a to	otal of sheets, as follows:						
sheets of the description, claims and/or drawings which have been amended and are the basis of this report and/or sheets containing rectifications authorized by this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions).								
T shoots which supersed	e earlier sheets, but which thi	is Authority considers contain an amendment that goes						
beyond the disclosure Supplemental Box.	beyond the disclosure in the international application as filed, as indicated in item 4 of Box No. I and the Supplemental Box.							
b. (sent to the International B	ureau only) a total of (indicate	type and number of electronic carrier(s)) , containing a er readable form only, as indicated in the Supplemental						
sequence listing and/or tab Box Relating to Sequence	Listing (see Section 802 of the	e Administrative Instructions).						
4. This report contains indications re	lating to the following items:							
☐ Box No. I Basis of the opin	nion							
☐ Box No. II Priority								
	ent of opinion with regard to n	ovelty, inventive step and industrial applicability						
☐ Box No. IV Lack of unity of								
☐ Box No. V Reasoned state								
☐ Box No. VI Certain docume	nts cited							
☐ Box No. VII Certain defects	in the international application	1						
☐ Box No. VIII Certain observa	tions on the international app	lication						
Date of submission of the demand	Date	of completion of this report						
19.10.2004		2.2005						
Name and mailing addrage of the international		orized Officer						
Name and malling address of the international preliminary examining authority:		Land Harden Palanten						
European Patent Office D-80298 Munich	Alla	utt, S						
7el. +49 89 2399 - 0 Tx: 523656 epmu d								
Fax: +49 89 2399 - 4465	Tele	phone No. +49 89 2399-7817						

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/050376

_	Box No. I	Basis of the report				
1.	With regard to the language , this report is based on the international application in the language in which it will filed, unless otherwise indicated under this item.					
	This report is based on translations from the original language into the following language, which is the language of a translation furnished for the purposes of:					
	 □ international search (under Rules 12.3 and 23.1(b)) □ publication of the international application (under Rule 12.4) □ international preliminary examination (under Rules 55.2 and/or 55.3) 					
2.	 With regard to the elements* of the international application, this report is based on (replacement sheets whi have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report): 					
	Description	, Pages				
	1-11	as originally filed				
	Claims, Nur	nbers				
1-22		as originally filed				
	Drawings, S	heets				
	1/4-4/4	as originally filed				
	□ a seque	ence listing and/or any related table(s) - see Supplemental Box Relating to Sequence Listing				
3.	☐ The an	nendments have resulted in the cancellation of:				
		description, pages claims, Nos.				
	☐ the	drawings, sheets/figs				
	☐ the ☐ any	sequence listing (specify): table(s) related to sequence listing (specify):				
4.	had not bee Supplement	port has been established as if (some of) the amendments annexed to this report and listed below an made, since they have been considered to go beyond the disclosure as filed, as indicated in the tal Box (Rule 70.2(c)).				
		description, pages claims, Nos.				
	☐ the	drawings, sheets/figs				
		sequence listing (specify): table(s) related to sequence listing (specify):				
	* If ite	em 4 applies, some or all of these sheets may be marked "superseded."				

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/050376

	Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability					
1.	The obv	ne questions whether the claimed invention appears to be novel, to involve an inventive step (to be non- ovious), or to be industrially applicable have not been examined in respect of:				
		the entire international applicat	entire international application,			
	\boxtimes	claims Nos. 9-15	s Nos. 9-15			
		because:				
	×	the said international application, or the said claims Nos. 9-15 (Industrial Applicability) relate to the following subject matter which does not require an international preliminary examination (specify):				
		see separate sheet		·		
		the description, claims or drawings (indicate particular elements below) or said claims Nos. are so unclear that no meaningful opinion could be formed (specify):				
		the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed.				
		no international search report has been established for the said claims Nos.				
		the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that:				
		the written form		has not been furnished		
				does not comply with the standard		
		the computer readable form		has not been furnished		
				does not comply with the standard		
		the tables related to the nucleon not comply with the technical re	tide a equire	and/or amino acid sequence listing, if in computer readable form only, do ements provided for in Annex C-bis of the Administrative Instructions.		
		See separate sheet for further of	detail	ds .		

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/050376

 C_{i}

Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N)

Yes: Claims

No: Claims

1-22

Inventive step (IS)

Yes: Claims

No: Claims

1-22

Industrial applicability (IA)

Yes: Claims

1-8,16-22

No: Claims

9-15

2. Citations and explanations (Rule 70.7):

see separate sheet

<u>ltem III</u>

٤

- 1. Claims 9-15 relate to subject-matter considered by this Authority to be covered by the provisions of Rule 67.1(iv) PCT. Consequently, no opinion will be formulated with respect to the industrial applicability of the subject-matter of these claims (Article 34(4)(a)(i) PCT).
- 2. The following documents are referred to in this communication; the numbering will be adhered to in the rest of the procedure:
 - D1: WO 03/011274 A (GLAXO GROUP LTD ;WARD PETER (GB); KNOWLES RICHARD GRAHAM (GB)) 13 February 2003 (2003-02-13)
 - D2: WO 02/069945 A (BOEHRINGER INGELHEIM PHARMA ;PIEPER MICHAEL PAUL (DE); PAIRET MICH) 12 September 2002 (2002-09-12)
 - D3: WO 02/096463 A (YEADON MICHAEL ;WATSON JOHN W (US); PFIZER (US); ARMSTRONG ROISIN) 5 December 2002 (2002-12-05)
 - D4: WO 02/096423 A (BOEHRINGER INGELHEIM PHARMA ;YEADON MICHAEL (GB); WATSON JOHN W (U) 5 December 2002 (2002-12-05)

The documents considered in the present processing are consecutively numbered D1-D4; this numbering results from the citations D1-D4 found in the Search Report (SR) of the corresponding PCT application. It will be adhered to in the rest of the procedure. The cited passage(s) for each citation will be considered unless otherwise specified.

Item V

<u>Novelty</u>

- 3. The subject matter of claims 1-22 are anticipated by prior art documents D1 and D2 and therefore do not fulfill the requirements of Art 33(2) PCT.
- D1 discloses a composition comprising tiotropium or a salt thereof in combination with a PDE4 inhibitor such as riflomilast for treating exacerbations associated with pulmonary disease e.g. COPD. The composition is administered via inhalation.
- D2 also discloses a composition comprising tiotropium, oxitropium and ipratropium salts in combination with a PDE4 inhibitor including riflomilast. The composition is administered by inhalation and is useful for treating inflammatory or obstructive pulmonary disorders.

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (SEPARATE SHEET)

International application No.

PCT/EP2004/050376

4. D3 discloses an inhalative combination of a PDE4 inhibitor and an anticholinergic agent other than tiotropium. Riflomilast was not specifically mentioned. D4 discloses an inhalative combination of a PDE4 inhibitor and an anticholinergic agent selected from e.g. a tiotropium derivative for treating COPD. Riflomilast was mentioned in the general context of PDE4 inhibitors but is not specifically mentioned in combination with riflomilast.

Further Remarks:

Industrial Applicability (Art 33(4) PCT).

5. For the assessment of the present claims 9-15 on the question whether they are industrially applicable, no unified criteria exist in the PCT Contracting States. The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognize as industrially applicable the subject-matter of claims to the use of a compound in medical treatment, but may allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a medicament for a new medical treatment.